2021 SISBID Unsupervised Lab

Genevera I. Allen, Yufeng Liu, Hui Shen, Camille Little

Data Description

gdat is Gene Expression Data, n = 445 patients x p = 353 genes

- Only 353 genes with somatic mutations from COSMIC are retained ## Data is Level III TCGA BRCA RNA-Sequencing gene expression data that have already been pre-processed according to the following steps:
- Reads normalized by RPKM
- Corrected for overdispersion by a log-transformation (1 + data)
- Short gene name labels are given as the column names

cdat is Clinical Data, n = 445 patients x q = 6 clinical features

- Subtype denotes 5 PAM50 subtypes including Basal-like, Luminal A, Luminal B, HER2-enriched, and Normal-like
- ER-Status estrogen-receptor status
- PR-Status progesterone-receptor status
- HER2-Status human epidermal growth factor receptor 2 status
- Node number of lymph nodes involved
- Metastasis indicator for whether the cancer has metastasized

Problems

Problem 1 - Dimension reduction

- 1a Apply PCA, NMF, ICA and MDS, UMAP, and tSNE to this dataset. Compare and contrast the results using these methods.
- 1b Relate the dimension reduction results with the clinical data. Is any clinical information reflected in the lower dimensional spaces?
- 1c Overall, which dimension reduction method do you recommend for this data set and why?

Problem 2 - Clustering

- 2a Apply various clustering algorithms such as K-means (explore different K), hierarchical clustering (explore different linkages), NMF, and biclustering. Compare the clustering results using these methods.
- 2b Relate the clustering results with the clinical data. Can the clustering algorithm recover some of the clinical information such as cancer subtypes?
- 2c Use Consensus Clustering to help Validate Clustering Results
- 2c Overall, which clustering method(s) do you recommend for this data set and why?

Problem 3 - Multiple comparisons

- 3a Identify important genes to differetiate ER postive and negative, PR postive and negative, HER2 postive and negative, metastasis status.
- 3b Try different procedures to adjust for multiple comparisons.
- 3c Examine the lists of genes identified using different methods for each clinical response. Which method is best? Why?

Problem 5 - Graphical models

5a - Use graphical models to explore interactions among genes. Are any of the well-connected genes related to patterns previously identified?

Problem 6 - Visulaization

- 6a Visualize this data using multiple approaches.
- 6b Prepare the "best" visual summary of this data.

Problem 7 - Exploratory Data Analysis Summary

- 7a What is the most interesting finding?
- 7b Is this finding consistent and stable?
- 7c Prepare a visual summary that best illustrates this interesting finding.

R scripts to help out with the BRCA case study Lab Don't peek at this if you want to practice coding on your own!!

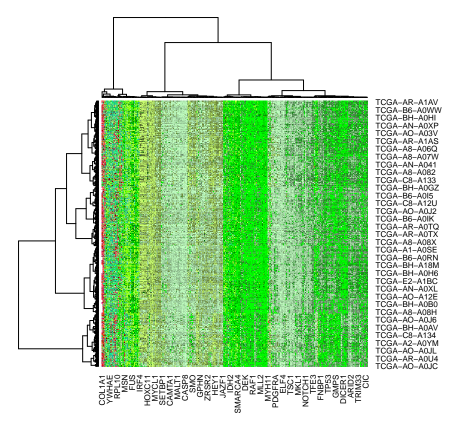
Load Data

```
load("UnsupL SISBID 2021.Rdata")
library(ggplot2)
## Warning: package 'ggplot2' was built under R version 3.6.2
library(ConsensusClusterPlus)
library(kknn)
library(GGally)
## Warning: package 'GGally' was built under R version 3.6.2
## Registered S3 method overwritten by 'GGally':
     method from
##
##
     +.gg
            ggplot2
library(umap)
## Warning: package 'umap' was built under R version 3.6.2
library(Rtsne)
library(igraph)
## Warning: package 'igraph' was built under R version 3.6.2
## Attaching package: 'igraph'
```

```
## The following objects are masked from 'package:stats':
##
##
       decompose, spectrum
## The following object is masked from 'package:base':
##
##
       union
library(huge)
## Warning: package 'huge' was built under R version 3.6.2
Explore Data
dim(gdat)
## [1] 445 353
dim(cdat)
## [1] 445
# clinical data
table(cdat$Subtype)
##
##
      Basal-like HER2-enriched
                                    Luminal A
                                                  Luminal B
                                                               Normal-like
##
              79
                                          200
                                                        106
table(cdat$ER)
##
##
                 Indeterminate
                                                   Negative
##
                                                         100
##
                 Not Performed Performed but Not Available
##
##
                      Positive
##
                            339
table(cdat$PR)
##
##
                 Indeterminate
                                                   Negative
##
##
                 Not Performed Performed but Not Available
##
##
                      Positive
                            291
table(cdat$HER2)
##
##
       Equivocal
                      Negative Not Available
                                                   Positive
                            370
                                                          65
table(cdat$Node)
##
##
    0 1
             2
                3
## 221 146 54 23
```

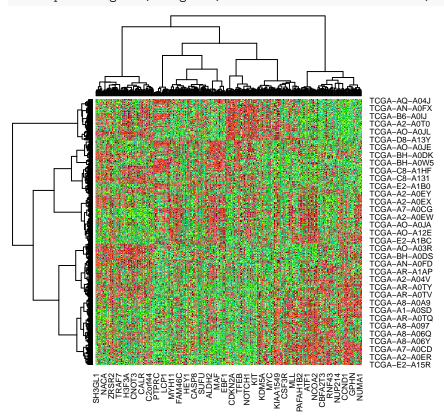
```
table(cdat$Metastasis)
##
##
    0
       1
## 427 11
table(cdat$ER,cdat$PR)
##
##
                                  Indeterminate Negative Not Performed
##
     Indeterminate
                                              0
                                                      1
##
     Negative
                                              1
                                                      93
                                                                     0
##
    Not Performed
                                             0
                                                       0
                                                                     2
    Performed but Not Available
##
                                             0
                                                       0
                                                                     0
##
    Positive
                                              2
                                                      53
##
##
                                 Performed but Not Available Positive
##
     Indeterminate
                                                            0
##
    Negative
                                                            0
                                                                     6
                                                            0
                                                                     0
##
     Not Performed
##
    Performed but Not Available
                                                            2
                                                                     0
##
    Positive
                                                            0
                                                                   284
#cluster heatmap - biclustering
#cluster heatmap - biclustering
aa = grep("grey",colors())
bb = grep("green",colors())
cc = grep("red",colors())
gcol2 = colors()[c(aa[1:2],bb[1:25],cc[1:50])]
Without scaling
```

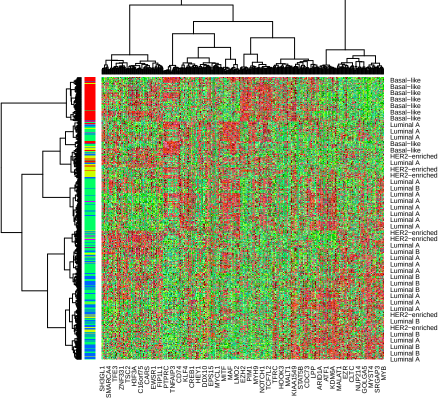
```
heatmap(gdat,col=gcol2,hclustfun=function(x)hclust(x,method="ward.D"))
```



With scaling

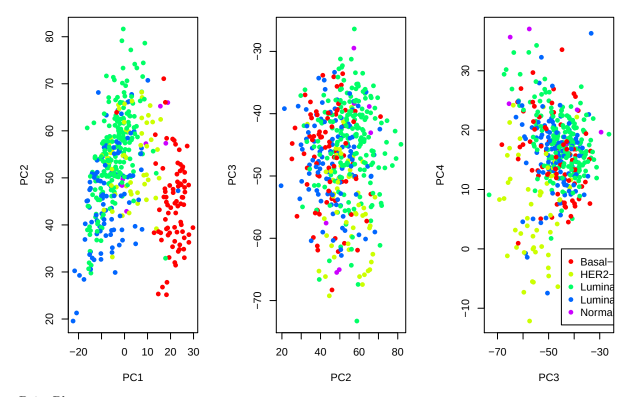
heatmap(scale(gdat),col=gcol2,hclustfun=function(x)hclust(x,method="ward.D"))





#Dimension Reduction

PCA

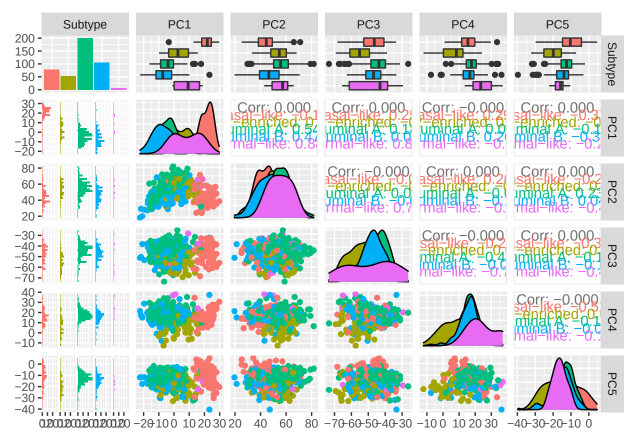


```
Pairs Plot

PC1<-as.matrix(Z[,1])
PC2<-as.matrix(Z[,2])
PC3<-as.matrix(Z[,3])
PC4<-as.matrix(Z[,4])
PC5<-as.matrix(Z[,5])

pc.df.cdat<-data.frame(Subtype = cdat$Subtype, PC1, PC2, PC3, PC4, PC5)
ggpairs(pc.df.cdat, mapping = aes(color = Subtype))

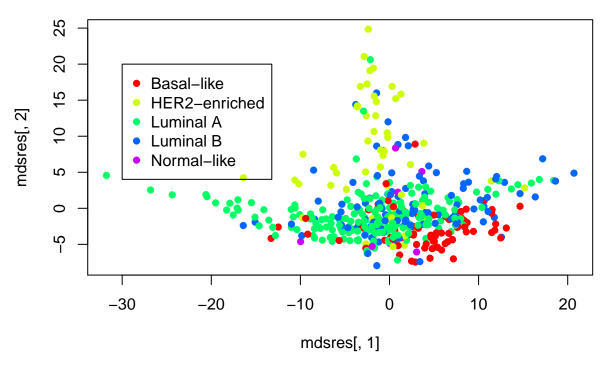
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.</pre>
```



MDS

```
Dmat = dist(gdat,method="maximum")
mdsres = cmdscale(Dmat,k=2)
plot(mdsres[,1],mdsres[,2],pch=16,col=Cols(cdat$Subtype), main = "Dimension Reduction MDS")
legend(-30,20,pch=16,col=rainbow(5),levels(cdat$Subtype))
```

Dimension Reduction MDS



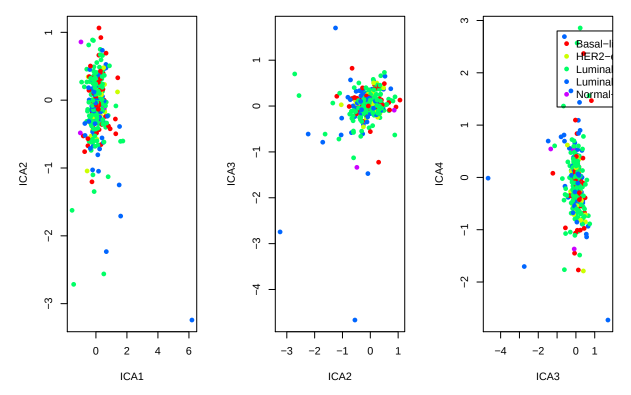
ICA

```
require("fastICA")

## Loading required package: fastICA

K = 4
icafit = fastICA(gdat,n.comp=K)

kk = 3
pclabs = c("ICA1","ICA2","ICA3","ICA4")
par(mfrow=c(1,kk))
for(i in 1:kk){
    j = i+1
    plot(icafit$A[i,],icafit$A[j,],pch=16,xlab=pclabs[i],ylab=pclabs[j],col=Cols(cdat$Subtype))
}
legend(-1,2.8,pch=16,col=rainbow(5),levels(cdat$Subtype))
```

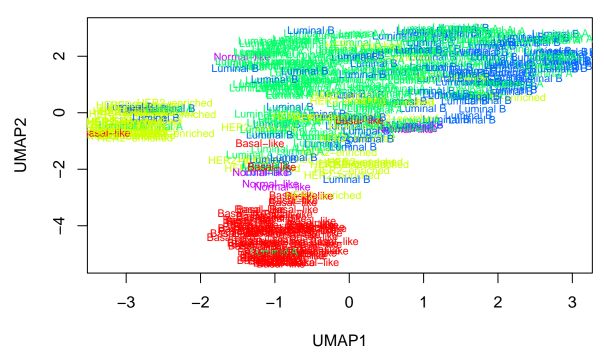


UMAP

```
gdat.umap = umap(gdat)
plot(gdat.umap$layout[,1], y = gdat.umap$layout[,2], type = "n", main = "UMAP", xlab = "UMAP1", ylab = "
text(gdat.umap$layout[,1], y = gdat.umap$layout[,2], type = "n", cdat$Subtype, col=Cols(cdat$Subtype), c
## Warning in text.default(gdat.umap$layout[, 1], y = gdat.umap$layout[, 2], :
```

graphical parameter "type" is obsolete

UMAP



#Clustering

K-means

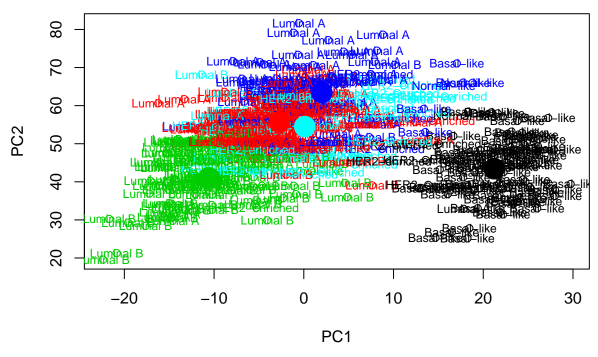
```
K = 5
km = kmeans(gdat,centers=K,nstart=25)
table(km$cluster,cdat$Subtype)
```

```
##
##
       Basal-like HER2-enriched Luminal A Luminal B Normal-like
##
     1
                74
                                 5
                                                        1
     2
                 0
                                 7
                                          112
                                                       26
##
                 0
                                 2
##
     3
                                           40
                                                       60
     4
                 5
                                 8
                                           44
                                                       12
                                                                     4
##
                                31
```

Plot Kmeans with labels

```
plot(Z[,1],Z[,2],col=km$cluster, main = "Plot Kmeans Clusters ", xlab = "PC1", ylab = "PC2")
text(Z[,1],Z[,2],cdat$Subtype,cex=.75,col=km$cluster)
cens = km$centers
points(cens%*%V[,1],cens%*%V[,2],col=1:K,pch=16,cex=3)
```

Plot Kmeans Clusters



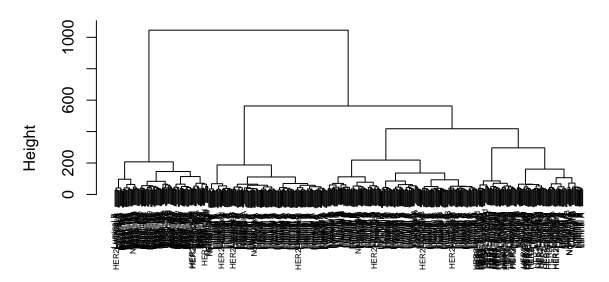
Hierarchical

```
#which linakge is the best?
#which distance metric is the best?

Dmat = dist(gdat)
com.hc = hclust(Dmat,method="ward.D")

plot(com.hc,labels=cdat$Subtype,cex=.5)
```

Cluster Dendrogram



Dmat hclust (*, "ward.D")

```
res.com = cutree(com.hc,5)
table(res.com,cdat$Subtype)
## res.com Basal-like HER2-enriched Luminal A Luminal B Normal-like
##
         1
                     1
                                    3
                                              95
                                                         11
         2
                     0
                                    4
                                              73
                                                         65
                                                                       1
##
                    75
##
                                    4
                                               5
                                                          4
                                                                       1
         4
                     0
                                   27
                                               3
                                                          7
##
                                                                       0
         5
                     3
                                   15
                                              24
                                                         19
```

Consensus Clustering with Hierarchical

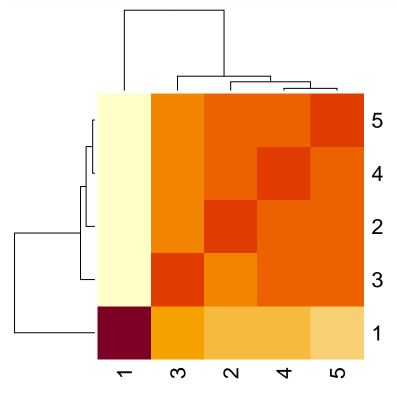
```
results = ConsensusClusterPlus(gdat,maxK=6,reps=500,pItem=0.8,pFeature=1,
clusterAlg="hc",distance="pearson",plot="png")
```

```
## end fraction
```

- ## clustered

Look at results for first 5 clusters

heatmap(results[[2]][["consensusMatrix"]][1:5,1:5])



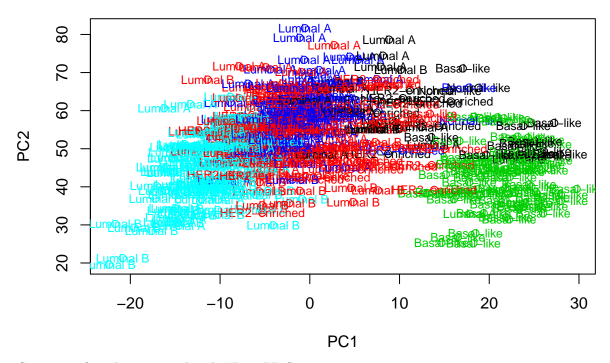
Spectral Clustering

```
K = 5
s_gdat = specClust(gdat, centers=K, nn = 7, method = "symmetric", gmax=NULL)
```

Visualize

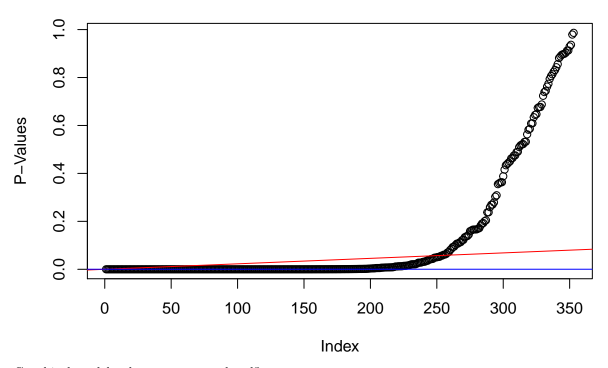
```
plot(Z[,1],Z[,2],col=s_gdat$cluster, main = "Visualize Spectral Clusters", xlab = "PC1", ylab = "PC2")
text(Z[,1],Z[,2],cdat$Subtype,cex=.75,col=s_gdat$cluster)
```

Visualize Spectral Clusters



Genes significantly associated with ER or PR Status, etc

```
x = gdat[cdat$ER=="Positive" | cdat$ER=="Negative",]
y.er = cdat$ER[cdat$ER=="Positive" | cdat$ER=="Negative"]
y.label = rep(1, length(y.er))
y.label[y.er == "Positive"]=2
ps = NULL
for(i in 1:ncol(gdat)) ps = c(ps,
t.test(x[y.label==1,i],x[y.label==2,i])$p.value)
fdrs.bh = p.adjust(ps, method="BH")
cat("Number of Tests significant with alpha=0.1 using Bonferroni correction:",
sum(ps<0.1/length(y.label)), fill=TRUE)</pre>
## Number of Tests significant with alpha=0.1 using Bonferroni correction: 165
cat("Number of Tests with FDR below 0.1:",
sum(fdrs.bh<0.1), fill=TRUE)</pre>
## Number of Tests with FDR below 0.1: 259
plot(sort(ps,decreasing=FALSE),ylab="P-Values")
#BH procedure
abline(a=0, b=0.1/length(y.label),col="red")
#Bonferroni
abline(a=0.1/length(y.label), b=0,col="blue")
```



Graphical models - how are genes related?

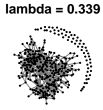
```
# use huge package
neth = huge(gdat,method="mb")
```

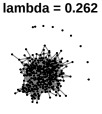
Conducting Meinshausen & Buhlmann graph estimation (mb)....done
plot(neth)

parsity vs. Regularization

1.0







Regularization Parameter

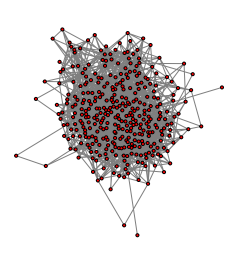
0.2

```
## stability selection with huge
net.s <- huge.select(neth, criterion="stars")</pre>
```

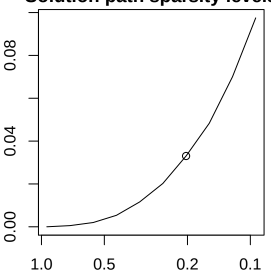
Conducting Subsampling....in progress:5% Conducting Subsampling....in progress:10% Conducting Subsamples...

```
## Model: Meinshausen & Buhlmann Graph Estimation (mb)
## selection criterion: stars
## Graph dimension: 353
## sparsity level 0.03304468
```

plot(net.s)

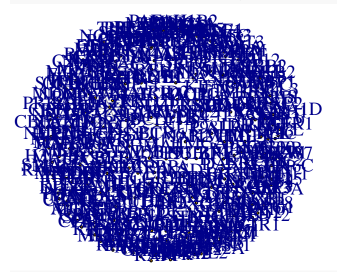


Solution path sparsity levels

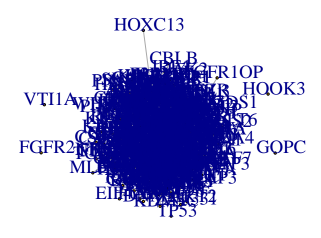


Regularization Parameter

```
#larger lambda
mat <- neth$path[[2]]
neti <- as.undirected(graph_from_adjacency_matrix(mat))
plot(neti,vertex.label=colnames(gdat),vertex.size=2,vertex.label.cex=1.2,vertex.label.dist=1,layout=lay</pre>
```



```
#smaller lambda
mat = neth$path[[6]]
neti = as.undirected(graph_from_adjacency_matrix(mat))
plot(neti,vertex.label=colnames(gdat),vertex.size=2,vertex.label.cex=1.2,vertex.label.dist=1,layout=lay
```



SS18